PATENT APPLICATION

IN THE UNITED STATES PATENT AND TRADEMARK OFFICE

In re Patent Application Serial No. 06/659,339

Filed: October 10, 1984

CANCELLATION OF PETITION FOR ACCESS TO PATENT APPLICATION PURSUANT TO 37 C.E.R. 1.14(1) AND (c)

Nutley, New Jersey 07110 July 24, 1996

Assistant Commissioner for Patents Washington, D.C. 20231

Attn: Special Program Law Office

Petition Information Crystal Park One Suite 520 FAX RECEIVED

JUL 1 Z 1996

PTITOUS OFFICE

Sir:

This paper is submitted to cancel the Petition For Access to Patent Application pursuant to 37 CFR 1.14(e)(1), which was mailed to the U.S. Patent and Trademark Office on June 11, 1996 by Catherine R. Smith. Enclosed is a copy of this Petition.

The undersigned attorney, on behalf of Ms. Catherine R. Smith who is on vacation, is requesting cancellation of this Petition. The undersigned attorney is the supervisor of Catherine R. Smith and together with Ms. Smith is employed in the Patent Law Department of Hoffmann-La Roche Inc., 240 Kingsland Street, Nutley, NJ 07110.

July 24, 1996

For: Mr. Hoffman

Deputy Assistant Commissioner's Office

Re: Abandoned Application Serial No. 06/659,339 - CHANG (USP 4,774,175)

I am FAXing herewith the cancellation of the Petition which was filed and directed to your department. Please let me know when this file will be available for me to copy same. Thanks. My number is 415-1224.

Bethy Boyed

CERTIFICATE OF MAILING (37 CFR 1.80)

I hereby certify that this paper (along with any paper referred	ed to as being transmitted therewith) in being deposited with the United in postage as first class mail in an envelope addressed to the: Assistant			
Commissioner for Patents, Washington, D.C. 20231.	(Print Name)			
•				
Date: 3-7 (1, 155)	(Signature)			

PATENT APPLICATION

IN THE UNITED STATES PATENT AND TRADEMARK OFFICE

In re Patent Application Serial No. 06/659,339

Filed: October 10, 1984

PETITION FOR ACCESS TO PATENT APPLICATION PURSUANT TO 37 C.F.R. L14(a) AND (e)

Nutley, New Jersey 07110 June 1996

Assistant Commissioner for Patents Washington, D.C. 20231

FAX RECEIVED JUL 7 & 1996

Attn: Special Program Law Office Petition Information

Crystal Park One Suite 520

PETITIOUS OFFICE

Sir:

Pursuant to 37 CFR 1.14(e)(1), the undersigned and appointees hereby petition for access to the above captioned patent application, to inspect and make copies thereof for the reasons set forth below.

The above patent application has been incorporated by reference in an issued U.S. Patent. U.S. Patent No. 4,774,175 issued September 27, 1988. A copy of the title page and page

> DEPOSIT ACCOUNT NO. 0843525

OUR ORDER NO.

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PAGE.04

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Serial No.

06/659.339

Filed:

October 10, 1984

encompassing columns 3 and 4 of that patent are attached hereto. In this regard attention is directed to column 4, lines 6 through 12, especially lines 8 through 9.

By incorporating by reference the above patent application into U.S. Patent No. 4,774,175, the right of confidentiality secured by 35 USC 122 has been waived with regard to the above application. (In re Yang, 177 USPQ 88 (Pat. Off. Sol. 1973); In re Gallo, 231 USPQ 496 (Commr. Pat. 1986; MPEP §103). Accordingly, the undersigned respectfully requests access to the captioned patent application as originally filed and to the file history thereof.

The undersigned submits that the requirements of 37 CFR 1.14(e)(1) have been met and respectfully requests that the present petition be granted.

The Assistant Commissioner of Patents and Trademarks is hereby authorized to charge the required petition fee of \$130.00, 37 CFR 1.17(I), and any other related fees to this petition or to credit any overpayment to Deposit Account No. 08-2525.

Respectfully submitted,

Attorney of Applicant(s)
Catherine R. Smith
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24060 Encl.

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United States Patent [19]

[11] Patent Number:

4,774,175

Chang et al.

[36]

Date of Patent:

Sep. 27, 1988

DMMUNOCHEMICAL METHODS FOR THE DETECTION OF ANTIBODY AGAINST HTLV-III,

[75] Inventors: Tm W. Chana, Paoli; Ikunoshin Kato, Exton; Pracab Chands; Nascy T. Chang, both of Paoli, all of Pa.

[73] Assignor: Ceatteor, Inc., Malvern, Pa.

[21] Appl. No.: 107,066

Mar. 1, 1985 [22] Filed:

Ex. CL. CLIQ L/70; COIN 33/544 [51] [52] U.S. Cl. 435/805; 436/318; 436/328; 436/331; 436/331; 436/811: 530/324

435/5, 7, 160, 805; 530/324; 436/518, 528, 531, 811 [58] Fichs of Search

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4.520,113 5/1983 Callo et al. . 4.629,783 12/1984 Consed _ \$30/324

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Primary Examiner-Christian M. Nucker Attorney, Agent, or Firm—Hamilton, Brook, Smith & Reynolds

ABSTRACT [57]

Gene segments of human T cell lymphotropic virus type III (HTLV-III) were expressed in E coli as peptides that are reactive with sera from patients with acquired inmoure deficiency syndrode (AIDS). Among recombinant peptides one designated HTLV-III polypeptide 121, contained 85 amino acid residues encoded by a gene segment in the est-lor region of the HTLV-III genome. The polypeptide is strongly reactive with AIDS patient sera. The peptide produced and purified as a fusion protein on a large scale. Solid phase immunosative employing this recombinant poptide as an immunosaborount can reliably and reproducibly detect antibodies in zera of putients with HTLV-IIII infection. In two representative serum panets, the stary detected the presence of antibodies in 120 of 121 zera from patients with AIDS or AIDS-related complex (ARC), and only in 1 of 92 normal controls. Based upon HTLV-IIII polypeptide 121 as immunotreanive agent, sensitive and specific immunoaxitys for HTLV-III infection have been developed. beca developed

15 Claims, 2 Drawing Streets

The polypeptide can be used in assays of various types including immunometric estays and anugen-sand-wich assays. A preferred type of assay is a solid phase immunometric double authority) assay, HTLV-III polymential 131 is inconstituted to strateging it to relief physics 121 is innobilized by attaching it to solid phase to thru an antigen immunoadsorbent. The im-munoadsorbent is used to adapt anti-HTLV-III and-body from a sample of the biological fluid. The adsorted ant-HTLV-III anabody is detected with in anti-(human IgO) anabody which is labeled radioistic- to pically, ensymmetricity, fluoremetrically or to other weys. This second antibody, directed generally against human IgG, binds to anti-HTLV-III antibody adsorbed human igG, hann to anti-HTLV-III authory attented to the immunoschorbest and produces a detectable nighal which can be evaluated as an indication of the 15 processor of anti-HTLV-III authory in the sample.

The in-munochemical stays employing HTLV-III polypopide 121 provide saveral advantages over those polypopide 121 provide saveral advantages over those human on the whole virus. A save based more HTLV-III

polypopoide 121 provide several savantages over those based on the whole virus Assays based open HTLV-III pulypopoide 121 efections to the send to grow large quantity of the inlegitous virus. This allowings the risk smoothed with this process. Additionally, sessy reagens based upon the HTLV-III satiges rather that the whole virus will help margate the real or perceived risk of contracting AIDS by technicisms who perform the 23 pages.

In performance, manys employing HTLV-LT polypeptide 121 are excellent, HLTV-III polypetide 121 are excellent, HLTV-III polypetide 121 are excellent, HTML polypetide 121 are excellent, For cample, in solid phase grays, background laint associated
with the immunosidantees is low. Further, the assays

are excellent highly sensitive and specific. Because are surprisingly highly sensions and specific. Because

ATC AGA TOC AAT AAT	GAG ATE TAE TET GET AAC	8368484	SASTERS	CTC GAT CAGE
AAT	TAC	AÇA	***	

ably exhibits for fewer episopes the polypeptide presse than the whole virus and consequently should be reac-tive with a smaller fraction of the antibudy against the virion, the high sensitivity and specificity was not experied. In the immensivity and specificity was not experied. In the immension ric many, the polypoptide
concreted the presence of anni-HTLV-III anniholy in
99% of arm of patients with A IPM and A O O Thomas 99% of zero of patients with AIDS and ARC. The very high specificity of the away suggests that HTLV-III polypepode 121 is derived from a highly amignic portion of the virtues and that antihody against the amignic evolud is virtually all instances of HTLV-III infection. These performance characteristics provide for highly american acrossing of blood and .ther bodily fluids for the presence of HTLV-III and for greater 15 precisions in the diagnosis of AIDS.

Because of the apparent arrong antigenicity of 99% of sera of passents with AIDS and ARC. The very

Because of the apparent strong antigenicity of HTLV-III polyappide 121, it could be used as a vaccine against the virus.

BRIEF DESCRIPTION OF THE DRAWINGS

PIG. 1 is a comparison between solid phase im-nanoadsorbents using recombinant HTLV-III antigenpolypesside 121, and inactivated, disrupted HTLV-III

FIG. 2 shows assay results on sera samples from patients with AIDS or ARC and from animal redividuats with immunoratiometre assays employing rection is

nant HTLV-III polypeptide 121 25 2 2016-phase imdeorbent.

BEST MODE OF CARRYING OUT THE HOLTHION

HTLV-III polypepude 121 was expressed and identified by a shorgun cloning procedure. This procedure is described in detail in U.S. patent application Ser. No. 659,339, which is incorporated by reference herein. For completeness, the procedure is outlined here and desended in further detail in the Exemplification Section

Closed HTLV-III DNA was broken into fragments Cloted HTLV-III DNA was broken into fragments of upproximately 500 base pairs in length and inserted into the "open reading frame" (ORF) cloning and expression vector pMR101. The inserted DNA was expressed in 8. and transformants as oripartice fusion proteins, constituing of an HTLV-III polypeptide fused to ACI constitue at its Naturalisan and between least confidence at ACI proctis at in N-terminal and beta-galacteridase at its C-terminal. About 300 closes were found to express to thermost. About AU closes were round to express bre-gulactuations activity indicating expression of the impartual fusion process. AIDS patient sera containing sub-HTLV-III antibodies were used to acreen for to-sion proteins that were immunoresceive. Among hreaty closes which produced proteins reserve with the AIDS sen, one close, designated close # 121, expresent a featon present which was immunorescrive with all AIDS periods are examined (24/24). The highly immunorescive protein produced by this close was relected for further study.

The HTLY-III DNA segment of clone 121 was exed from pMR 100 and sequenced by the Sanger techaigue. The auxicoade sequence is as follows:

CAG	CAG	1835	ACT ACT	CE8533	STE	100 CCT	TOG
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Based upon the DNA sequence, the putative amino acid sequence of the HTLV-III polypeptide could be

FASAR	A 1 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2	2985995	Gly Ala Len Len Ala	Glo lie is to a lie is to a lie	CA CA	ATE IN THE SET	Try Tar Ass	Lev Giy A by Lyr	1,5 5 as 4 5 5 5 5 5 5 5 5 5 5 5 5 5 5 5 5 5 5
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pMR 100 prepartice fusion protein synthesized by close 121 was difficult to purify in sufficient quantity for sera screening because the expression level was low (approximately 1.0% of total cellular protein) and the protein was insoluble in conventional extractive buffer protein was insoluble in conventional extractive buffer (probably due to the existence of 23 half-aystind francues). In order to enhance expression the HTLV-III polyperpide in E coll the HTLV III segment of clone 121 was closed into a beta-glucuronidase expression vector. E coll transformed with the recombinant vector capterised a 15 Kd fusion protein with short fusion participal at both ends (41 amino acids of E cole the 83 amino and colling and the 12 amino acids of E cole the 83 amino acids of E cole the 84 amino acids of E cole acid residues encoding by HTLV-III polypeptide ind 13 amino acid residues encoded by a multiple close as